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- (54) A coll coating paint composition.

A coil coating composition consisting essentially of (A) a binder component comprising a mixture of a linear polyester resin and a branched polyester resin, having a hydroxyl value of 40 to 90, and (B) a crosslinking component selected from melamine resin, at least part of which is methyl methylolated melamine, or isocyanate compound. Said polyester resins are herein defined.

The coating composition is useful in a coil coating and results a coating with excellent properties in respect of hardness, bending processability, fastness to boiling water, weather resistance, chemical resistance, marker stain resistance and the like.

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A coll coating paint composition

The present invention relates to a coil coating paint composition. More specifically, the invention concerns a high solids coating composition for coil coating use, capable of forming a coating with excellent properties in respect of hardness, bending processability, fastness to boiling water, weather resistance, chemical resistance, marker stain resistance and the like.

As a coating composition to be used in a coil coating of steel plate and the like, various properties including excellent weather resistance, chemical resistance, water resistance, coin scratch resistance, gloss, hardness, and bending properties of the coating, as well as the capability of formulating a high solids paint.

In such application, a polyester resin is a promising material because of resulting a coating with excellent weather resistance, chemical resistance, hardness and the like. However, this material has a serious drawback of lacking in structural flexibility. As an attempt for giving strechability and flexibility to a polyester resin, has been proposed a technique wherein a comparatively long methylene chain is included in either of dicarboxylic acio or diol component or in lactone to be reacted with polyester resin, thereby introducing the same into the polyester's molecular structure. However, haroness and bending properties are, in principle, contrary to each other, and an improvement in flexibility is always linked with the decrease in hardness. Thus, it is quite difficult to attain the well balanced flexibility and hardness in a single polyester (resin.

It has also been proposed to use a mixture of soft polyester and hard polyester as a binder resin (Japanese Patent Application Kokai No. 111864/83, ibid 111865/83). In these attemps, while maintaining hardness, water resistance and chemical resistance each in an appropriate level, it was indeed succeeded in having an improved flexibility, rubber-like elasticity and the like by the control of methylene chain content of the resinous composition, but it was only settled to the mere satisfaction of the desired properties of the coating on soft materials as fender, bumper and the like. The proposed coating compositions were of quite dissatisfaction when used as a coil coating of steel plate especially with respect to hardness, bending properties and the like.

Recently, a very attractive lacquer type coil coating paint composition has been proposed in Japanese Patent Application Kokai No. 210975/84, in which a linear polyester resin having a number average molecular weight of 15,000 to 50,000 and a linear or branched polyester resin having a number average molecular weight of 2,000 to 10,000 are mixed in a weight ratio of 80:20 to 20:80 and used as a resinous binder. The said composition is said to give a coating with well balanced hardness and bending processability.

However, since the coating composition does not include any crosslinking agents, the thus formed coating is insufficient in weather resistance, fastness to boiling water, chemical resistance and the like. Even if a crosslinking agent is tried to be blended with said coating composition, an additional problem of poor compatibility would be inevitably arisen from the high molecular weight of the polyester resin used.

Therefore, the application of said lacker type paint composition is in fact confined in limited areas.

An object of the present invention is therefore to provide a polyester binder which is compatible with a crosslinking agent. An additional object of the invention is to provide a coating composition for coil coating use, which will give the coating with well balanced hardness and bending processabilities, as well as the characteristic properties possessed by polyester resin, and capable of being formulated as a high solids paint.

The inventors, as the results of extensive studies, have found that if a melamine resin at least part of which is methyl methylolated melamine (e.g. methylol melamine methyl ether alone or a mixture of methylol melamine methyl ether and methylol melamine butyl ether) or an isocyanate compound is selected as a crosslinking agent and a combination of low molecular weight (up to number average molecular weight 5000) linear polyester and a branched polyester is used as binder resins, a compatible blend can be formulated and a high solids coating composition capable of resulting a coating with well balanced hardness and bending processability can be obtained. On the basis of these findings, the present invention has been made.

Thus, according to the invention, is provided a coating composition for coil coating use consisting essentially of

(a) a binder component comprising a mixture of a linear polyester resin and a branched polyester resin, having a hydroxyl value of 40 to 90, said linear polyester resin being composed of polybasic acid component and polyhydric alcohol component consisting of trihydric and/or polyhydric alcohol (0 to 10wt% of the solid matter) and dihydric alcohol and having 1.8 to 2.2 hydroxyl groups per molecule and a number

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average molecular weight of 1000 to 5000, and said branched polyester restn being composed of polybasic acid component and polyhydric alcohol component consisting of trihydric and/or tetrahydric alcohol (3 to 40 weight% of the solid matter) and dihydric alcohol, and having 2.3 to 4.5 hydroxyl groups per molecule, and

(B) a crosslinking component consisting of melamine resin, at least part of which is a methyl methylolated melamine, or isocyanate compound.

The first polyester resin used in the present invention is an oil-free linear polyester resin composed of polybasic acid component and polyhydric alcohol component, the polyhydric alcohol component consisting of substantial amount of dihydric alcohol and a small amount (0 to 10 weight % of the solid matter) of trihydric and/or tetrahydric alcohol, having 1.8 to 2.2 hydroxyl groups per molecule and a number average molecular weight 1000 to 5000. By the employment of this type of polyester resin, entanglement of resinous molecules and physical crosslinking are of use in giving improved hardness, weather resistance and fastness to boiling water of the coating, and easy unraveling properties of the entangled molecules, sliding properties and stress relaxation are believed to be of use in giving improved bending properties and low temperature impact strength. When the hydroxyl groups are less than 1.8 per molecule, there remains a considerable number of unreacted chains, which will cause the decrease in weather resistance of the coating, whereas if the hydroxyl groups exceed over 2.2 per molecule, there will cause the decrease in sliding properties and hence the desired bending properties.

The number average molecular weight of said linear polyester resin should be in a range of 1000 to 5000, and preferably 2000 to 4000. This is because, in the combination of said resin with the specified crosslinking agent and branched polyester resin, if the number average molecular weight of said linear polyester is less than 1000, it is unable to expect the desired bending processability, and if it exceeds over 5000, it is unable to have a high solids coating composition and the desired compatibility with the selected crosslinking agent.

The second polyester resin used in the present invention is a branched, oil-free polyester resin composed of polybasic acid component and polyhydric alcohol component consisting of trihydric and/or tetrahydric alcohol, the hydroxyl groups number per molecule being 2.3 to 4.5. By the selective use of this type of branched polyester resin, hardness, weather resistance and fastness to boiling water of the coating are greatly improved because of the increase in chemical crosslinking between the functional groups contained therein.

If the hydroxyl group number is less than 2.3, it is unable to obtain the desired hardness and weather resistance of the coating, whereas the preparation of branched polyester resin having more than 4.5 hydroxyl groups per molecule is very difficult to do.

In the present invention, the abovementioned linear polyester resin and brancheo polyester resin are mixed together so as to give a hydroxyl value of 40 to 90. This is because, if the hydroxyl value of the mixed resin is less than 40, the formed coating is too soft to the objects of the invention, and if it exceeds over 90, the coating is too hard.

As to the number average molecular weight of said branched polyester resin, it is not so critical to the objects of the invention and is approxiately selected in a common range of about 800 to 4000.

The present branched polyester resin may be modified with a lactone of the formula:

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wherein n is an integer of 4 to 10, thereby giving softness to the branched polyester resin and increasing crosslinking density of the cured coating, if desired. Said lactone may be used in an amount of 0 to 40 weight% of the resin solid. Further more, in the present invention, either or both of the abovementioned linear and branched polyester resins may be modified with an alkyleneimine as, for example, N-(2-hydroxy ethyl) ethyleneimine (HEEI) to improve pigment dispersion properties of the resin composition.

As already stated, the most characteristic feature of the invention resides in the point that the abovementioned resin composition is used as a binder component together with a crosslinking agent selected from the group consisting of a melamine resin at least part of which is methyl methylotated melamine, and an isocyanate compound.

Contrary to our expectations, number average molecular weight of the linear polyester resin to be blended with a branched polyester resin could be markedly lowered when the abovesaid particular compound was selectively used as a crosslinking agent.

At present, the exact reasons have not been fully understood, but the following are presumed to be one of the grounds thereof.

That is, when said crosslinking agent is selected, differing from the case with butylated melamine, the self condensation of the crosslinking agent is prevented and the resin and the crosslinking agent are reacted with each other in alternative order to give a soft, cured coating.

Lowering of the molecular weight of the polyester resin contributes to the improvement in the compatibility of said resin with a crosslinking agent and to the formation of a high solids coating composition. Employment of said polymer blend can result well balancec properties of hardness and bending processability, and employment of said crosslinking agent far improvements in weather resistance, fastness to boiling water, chemical resistance and marker stain resistance of the coating. Thus, the present composition is very useful as a resinous vehicle of a coil coating paint composition.

As to the blending ratio of the abovementioned polyester resins, it is selected in a wider range depending on the application desirec. The polyester blend is preferably compounded with the abovementioned crosslinking agent in a weight ratio (as solid) of 5/95 to 50/50, most preferably 20/80 to 35/65.

The present coating composition is usually applied on a steel plate or the like by coil coating means and baked, in most cases, at 180 to 250°C for 30 seconds to 2 minutes to give a coating with excellent properties with respect to hardness, bending properties, weather resistance, gloss, coin scratch resistance, fastness to boiling water, chemical resistance and marker stain resistance.

The invention shall be now more fully explained in the following Examples. Unless otherwise being stated, all parts and % are by weight.

Synthetic Example 1

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Preparation of linear polyester resin (A)

Into a 2 liters reaction vessel fitted with a stirrer, a fractionating column, a nitrogen gas inlet tube and a thermometer, were placed 140.9 parts of terephthalic acid, 495.5 parts of adipic acid, 135.9 parts of ethyleneglycol, 227.8 parts of neopentylglycol and 2 parts of dibutyl tin oxide and the mixture was heated. At the stage when it was reached to a stirrable condition, stirring was commenced to start and the temperature was raised to 240°C. At that time, from 160°C to 240°C, the temperature was raised at a constant speed in 3 hours. The raction water formed was continuously distilled off through the fractionating column out of the system. When the temperature was reached to 240°C, the mixture was maintained at said temperature and stirred for 1 hour. Thereafter, the fractionating column was stopped from working, xylene was added and the mixture was further reacted under refluxing conditions. When the acid value of the reaction product was reached to about 2.0, the reaction was overed and the mixture was allowed to cool. After cooling, 214 parts of xylene and 214 parts of Cellosolve acetate were added to obtain an oil-free polyester resin solution A-O. Analysis showed that it had a solid content of 70.4%, a viscosity (Gardner bubble viscometer, 25°C) of Y and contained polyester resin having a number average molecular weight of 5.000, a hydroxyl value of 21, an average number of hydroxyl groups per molecule of 1.8 and an acid value of 2.1.

Synthetic Examples 2 to 5

Following the procedures of Synthetic Example 1 but using the materials shown in Table 1, various linear polyester resins were prepared. However, in these Examples, a light modification was made as follows. That is, in the cases of the resins A- \oplus and A- \bigcirc , the condensation reactions were stopped at the stage when the acid value (AV/NV) was reached to 7.0 and 5.0, respectively, and the reaction mixture was allowed to cool to 90°C. While keeping the temperature at 90°C, 7.0 parts (in the case of A- \bigcirc) and 4.3 parts (in the case of A- \bigcirc) of N-(2-hydroxyethyl)ethyleneimine (HEEI) were added and the mixtures were reacted for 1 hour and then diluted with a mixture of 269 parts of xylene and 269 parts of Cellosolve acetate, respectively. The characteristics of these resin varnishes are shown in Table 1.

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Synthetic Example 6

Preparation of branched polyester resin (B)

Into a similar reaction vessel as used in Synthetic Example 1, were placed 373.9 parts of isophthalic acid, 85.6 parts of hexahydrophthalic anhydride, 94.6 parts of 1,6-hexahediol, 81.1 parts of neopentylglycol, 103.6 parts of trimethylol propane, 229.7 parts of Cardula E-10, and 2 parts of dibutyl tin oxide and the mixture was heated. At the stage when it was reached to a stirrable condition, stirring was commenced to start and temperature was raised to 230°C. At that time, from 195°C to 230°C, temperature was raised at a constant speed in 2 hours. Thereafter, the reaction was continued as in Synthetic Example 1 until the acid value of the reaction product was reached to about 3.0. After cooling to 150°C, the mixture was added with 31.5 parts of phthalic anhydride and reacted to obtain an oil-free polyester resin solution. The characteristics of thus obtained varnish are shown in Table 1.

Synthetic Example 7

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Into a similar reaction vessel as used in Synthetic Example 1, were placed 274.0 parts of isophthalic acid, 255 parts of hexahydrophthalic anhydride, 178.3 parts of 1,6-hexahediol, 157.1 parts of neopentyl-glycol, 61.5 parts of trimethylol propane, 73.8 parts of caprolactam and 2.0 parts of dibutyl tin oxide and the mixture was reacted as in Synthetic Example 1 until the resinous acid value was reached to about 15. After cooling, the mixture was added with 269 parts of xylene and 269 parts of Cellosolve acetate to obtain a resin varnish. The characteristics of thus obtained varnish are shown in Table 1.

Synthetic Example 8

Repeating the same procedures as stated in Synthetic Example 7 but using the materials shown in Table 1, an oil-free polyester resin varnish was prepared, whose characteristics were also shown in Table 1.

Synthetic Example 9

Into a similar reaction vessel as used in Synthetic Example 1, were placed 288.1 parts of isophthalic acid, 67.1 parts of hexahydro phthalic anhydride, 94.6 parts of neopentylglycol, 124.9 parts of trimethylol-propane, 182.9 parts of Cardula E-10 and 207.2 parts of ε-caprolactam and the mixture was reacted as in Synthetic Example 1. As soon as the resinous acid value was reached to about 7.0, the reaction was stopped and the reaction mixture was allowed to cool to 140°C. 35.3 parts of phthalic anhydride were added and the mixture was further reacted to obtain an oil-free polyester resin varnish. The characteristics of thus obtained varnish are shown in Table 1.

Synthetic Examples 10 to 12

Repeating the same procedures as stated in Synthetic Example 1 but using the materials shown in Table 1, branched polyester resin varnishes were prepared. The characteristics of thus obtained varnishes are shown in Table 1.

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Synthetic Example 13

Preparation of Comparative resin varnish C- (13)

Into 2 liters reaction vessel which was similar to that of Synthetic Example 1 and was connected to a vacuum pump, were placed 244.9 parts of terephthalic acid, 244.9 parts of isophthalic acid, 298.3 parts of sebacic acid, 138.4 parts of ethyleneglycol, 232.4 parts of neopentylglycol, 2 parts of bibutyl tin oxide and 2 parts of antimony trioxide and the mixture was reacted at 170 to 250°C. After completion of the esterification reaction, polycondensation reaction was carried out under reduced pressure of 0.5 to 1.0 mmHg to obtain a high molecular weight linear polyester resin. The characteristics of thus obtained resin 6> 20000 also grope als in 01028 varnish are shown in Table 1.

Synthetic Example 14

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Preparation of Comparative resin varnish C-(14)

Repeating the same procedures as stated in Synthetic Example 1 but using the materials shown in Table 1, a low molecular weight linear polyester resin varnish was prepared. The characteristics of said resin varnish are shown in Table 1. 5 800 also Received in O1028

Synthetic Example 15

Preparation of Comparative resin varnish C-

Repeating the same procedures as stated in Synthetic Example 1 but using the materials shown in Table 1, a linear polyester resin varnish was prepared. The characteristics of thus obtained varnish are L> 04 2=28 -> Aleirer als en 01028 shown in Table 1.

Synthetic Examples 16 and 17

Paration of Comparative resin varnishes C-(1) and C-(1)

Repeating the same procedures as stated in Synthetic Example 1 but using the materials shown in the 1, branched polyester reisn varnishes were prepared. The characteristics of the same procedures are stated in Synthetic Example 1 but using the materials shown in Preparation of Comparative resin varnishes C-1 and C-1

Table 1, branched polyester reisn varnishes were prepared. The characteristics of these varnished are shown in Table 1.

In the following Example, a number average molecular weight of a polyester resin is calculated by the equation: (as prescribed in "A guide to synthetic resins for coating use" written by Kyozo Kltaoka, published by High Molecular Publishing Associate, page 101)

$$M_n = \frac{W}{(ma + mb) - (1 - \frac{\alpha}{100}) ea_2 - ea_1 + \frac{W - AV}{56100}}$$

α: intramolecular reaction rate % (in case of phthalic anhydride ... 4.0)

W: weight of final resin

ma: number of moles of acid in the final resin mb: number of moles of alcohol charged

eaz: equivalent of acid charged

eat: equivalent of monobaric acid charged

AV: acid value of final resin

Example 1

A coil coating composition was prepared by kneading the following mixture in a ball mill for 24 hours. The oil-free polyester resins A- ① and B- ⑥ used are shown in Synthetic Examples 1 and 6 respectively. Formulation:

	70.4% linear polyester resin A-①	ı	71.4	parts
10	65.0% branched polyester resin B-	6	76.9	
	80% methyl methylolated melamine	1)	37.5	
•	titanium oxide	2)	100.0	
15	p-toluene sulfonic acid solution	3)	6.5	
•	Modaflow 4)		0.3	
20	n-butanol		15.0	
	•	Total	307.6	•

1) Sumimal M-40S, non-volatile content 80%, trade mark, Sumitomo Chem. Co.

2) Titanium CR-91, trade mark, Ishihara Sangyo

3) 20 wt% benzylalcohol solution>

4) trade mark, Monsanto Cem. Co.

Thus obtained composition was added with a dilution thinner: 1:1 (by weight) = Solvesso 150 : Isophorone

to give a diluted composition having a viscosity of 120 seconds (Ford Cup #4, at 20°C).

The diluted coating composition was then applied on a galvanized steel having a thickness of 0.3mm (JIS-G3302) previously treated with a zinc phosphate solution, to give a wet film thickness of 10 to 15μ and the coated steel plate was baked at 230°C for 40 seconds.

The following tests were carried out with the coating composition and with the coated plate.

1) non-volatile content (%) of the coating composition

A coating composition was diluted with a mixed dilution thinner (isophorone/Solvesso #150 = 1/1 wt/wt) to a viscosity (#4 Ford cup. at 25°C) of 120 seconds. An amount of said composition was taken and weighed, and after being subjected to a drying at 105°C for 3 hours, the residual amount was again weighed and non-volatile content of said composition was calculated therefrom.

2) gloss

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60° reflextion factor was ditermined by using Murakami type GM-3H glossmeter.

3) Pencil hardness

Pencil hardness was determined by using a high quality pencil prescribed by JIS-S-6006 and following a standardized method of JIS-K-5400

4) bending properties

A coated steel was folced at 180° and cracking behavior at the bended portion was observed.

2T, for example, stands for no cracking when two coated plates having the same thickness are bended together.

5) fastness to boiling water

A coated plate was dipped in a boiling water for 8 hours, and the coating was then observed.

O... good ×... no good (generation of blister)

6) chemical resistance

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0.5cc of 10% aq. H₂SO₄ solution was dropped on a coated surface and the plate was maintained at 20°C and under 75% relative humidity (RH) condition for 24 hours. Thereafter, the coating was washed with water and chemical resistance of the coating was determined.

O... good ×... no good

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7) marker stain resistance

A red colored oil ink was applied on the coating (in 0.5x2cm area) and after leaving it at 20°C for 24 hours, the inked area was scrubbed with a gauze impregnated with methanol and observed the degree of stain

O... good × ... no good

30 8) weather resistance

Gloss retention (%) was measured after 2000 hours' irradiation in Sunshine weather-O-meter. The test results are shown in Table 2.

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Example 2 and 3

Using the linear polyester resins A- ② and A-③ and the branched polyester resins B- ⑤ and B-⑤, the same procedures as stated in Example 1 were repeated to obtain coating compositions. The respective coating composition was diluted, applied on the similar galvanized steel, baked and tested as in Example 1.

The test results are shown in Table 2.

Example 4

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A coil coating composition was prepared by using the oil-free polyester resin A- ② and B- ③ obtained in Synthetic Examples 4 and 9, and following the dispersion formulation (1) and solution formulation (2) hereinunder stated.

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Dispersion formulation (1) 65.2% linear polyester resin A-4 46.2 parts 100.0 titanium oxide 15.0 n-butanol Total 161.2 10 Solution formulation (2) Dispersion paste 15 161.2 parts (of dispersion formulation (1)) 65.0% branched polyester resin B-9 37.5 80% methyl methylolated melamine p-toluene sulfonic acid solution 6.5 0.3 Modaflow 25 Total 273.0

Thus obtained coating composition was diluted, applied on a galvanized steel previously treated with a zinc phosphate solution and baked as in Example 1.

The same tests as stated in Example 1 were carried out and the test results were shown in Table 2.

Examples 5 and 6

Using the oil-free polyester resins A- and B- obtained in Synthetic Examples 4 and 10 in Example 5, and the oil-free polyester resins A- and B- obtained in Synthetic Examples 5 and 8 in Example 6, the same procedures as stated in Example 4 were repeated to obtain coating compositions. They were diluted, applied on galvanized steel plates previously treated with a zinc phosphate solution, baked and tested as in Example 1. The test results are shown in Table 2.

Examples 7 and 8

Using the oil-free polyester resins A- ② and B- ① obtained in Synthetic Examples 2 and 11 in Example 7 and the oil-free polyester resins A- ③ and B- ② obtained in Synthetic Examples 3 and 12 in Example 8, together with an amount of methyl methylolated melamine shown in Table 2, the same procedures as stated in Example 1 were repeated to obtain coating compositions. They were diluted, applied on galvanized steel plates previously treated with a zinc phosphate solution, baked and tested as in Example 1. The test results are shown in Table 2.

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Example 9

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A coil coating composition was prepared by using the oll-free polyester resins A- ① and B-⑥ obtained in Synthetic Examples 1 and 6 and Following the prescription metnioned below:

	70.4% linear polyester resin A-①	85.2 parts
	65.0% branched polyester resin B-6	61.5
10.	80% methyl methylolated melamine	26.3
	60% butyl methylolated melamine 1)	15
15	titanium oxide	100
	p-toluene suflomic acid soltion	3.0
	Modaflow	0.3
20	n-butanol	15.0
	Total	306.3

1) ... Super Beckamine J-820-60, trade mark, Dainippon Ink
Chem. Ind. Ltd., non-volatile content 60%

Thus obtained coating composition was diluted, applied on a galvanized steel plate previously treated with a zinc phosphate solution, baked and tested as in Example 1. The test results are shown in Table 2.

Example 10

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Using the oil-free polyester resins A- ② and B- ⑦ obtained in Synthetic Examples 2 and 7 and the melamine resins shown in Example 9, the same procedures as stated in Example 9 were repeated to obtain a coating composition. The composition was diluted, applied on a galvanizeo steel plate previously treated with a zinc phosphate solution, baked and tested as in Example 1.

The test results are shown in Table 2.

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Example 11

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A coil coating composition was prepared by using the oil-free polyester resins A- 2 and B- 1 obtained in Synthetic Examples 2 and 11 and following the prescriptions shown below.

Dispersion formulation (1)

70.2% linear polyester resin A-② 42.7 parts titanium oxide 100.0

Total 142.7

Solution formulation (2)

Dispersion paste

(of disperion formulation (1)) 142.7 parts
70.2% linear polyester resin A-2 71.2
65.2% branched polyester resin B-1 30.7
Sumidule N-75 1) 16.0
dibutyl tin dilaurate 0.02
Modaflow 0.5

Total 261.1

1)... 75% aliphatic polyisocyanate in 1:1 Cellosolve acetate/xylene, trade mark, Sumitomo Bayer Urethane K.K.

Thus obtained coating composition was diluted, applied on a galvanized steel plate previously treated with a zinc phosphate solution, baked and tested as in Example 1. The test results are shown in Table 2.

Comparative Examples 1 to 9

Comparative coating compositions 1 to 9 were prepared by using the same procedures as stated in Example 1 and the materials shown in Table 2. They were diluted, applied on a galvanized steel plate previously treated with a zinc phosphate solution, baked and tested as in Example 1. The test results are also shown in Table 2.

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Table 1

linear type polyester resin

•					
Synthetic Example	1	2	3	4	5
oil-free polyester resin	A-(1)	A-2	A-3)	A-4	A-(5)
Composition and characteristics					
resin raw materials					(
1. terephthalic acid	140.9	373.0	137.1		`
2. isophthalic acid				154.7	195.7
3. phthalic anhydride			•	341.5	220.7
4. adipic acid	495.5		482.5	85.0.	122.3
5. azelaic acid		147.0		٠	
6. sebacic acid		147.0		-	
7. hexahydro phthalic anhydride					
8. ethyleneglycol	135.9	124.0	91.1		
9. 1,4-butanediol	•		132.1		
10. 1,6-hexanediol					242.0
11. neopentyl glycol	227.8	209.0	153.2	348.8	176.0
12. trimethylol propane	•		4.0	77 • 8·	43.3
13. pentaerythritol			٠.		
14. Cardura E-10					
15. E-caprolactone(n=5)			•		
16. phthalic anhydride					•
(for later charge)		•			•
17. 2-hydroxyethyl					
ethyleneimine (HEEI)	:			7.0	4.3

Table 1 (continued)

linear type polyester resin

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	c.	₽.	Α.	lo.	o	<u>a</u>	ο·	٥.	ت •	i PC	Con	oil	Syn
10	acid va	viscosi	non-vol	characteristics	<pre>6-caprolactone (wt%)</pre>	trivale	average	OH value	number average	resin spe	position	oil-free po	Synthetic Example
15	acid value (AV/NV)	ty (25°С	non-volatile content	ristics	lactone	trivalent/tetravalent	OH numbe	•	werage n	specifications	and cha	polyester	xample
20	NV)	viscosity (25°C Gardner)	ntent (%)	of resin	(wt%)	valent alo	average OH number per molecule		molecular weight	ons	Composition and characteristics	resin	÷
25						alcohol (wt%)	lecule		weight		ics		
30				-		*			UII				
	2.1	Y	70.4		i	0	1.8	21	5,000			A-O	
35	2.0	$^{2}1$	70.2		.	С	1.9	35	3,000			A-2	2
40	2.0	ĸ	69.8		· i	0.4	2.0	4 5	2,500			A-(3)	ω
45	2.1	ΥX	65.2		1	7.8	2.2	62	2,000			A-4	4
50	2.2	rŝ	65.0		1	<u>.</u> ω	2.0	94	1,200			A-(5)	un
	•					•							

Table 1 (continued)

branched type polyester resin

Synthetic Example	6 ·	7 .	8	9	10
oil-free polyester resin	B-6	B-(7)	B - (8)	в-9	B-10
Composition and characteristics					
resin raw materials					€
l. terephthalic acid					• *.
2. isophthalic acid	373.9	274.0	455.5	288.1	205.7
3. phthalic anhydride					
4. adipic acid					120.6
5. azelaic acid	,		•		
6. sebacic acid					
7. hexahydro phthalic anhydride	85.6	255.2		67.1	
8. ethyleneglycol					
9. 1,4-butanediol					
10. 1,6-hexanediol	94.6	.178.3	119.9		(
11. neopentyl glycol	81.1	157.1	160.0	94.6	156.6
12. trimethylol propane	103.6	61.5	52.1	124.9	116.8
13. pentaerythritol		-			
14. Cardura E-10	229.7			182.9	
15. &-caprolactone(n=5)		73.8	212.5	207.2	400.0
16. phthalic anhydride	31.5			35.3	
(for later charge)					
17. 2-hydroxyethyl			••		
ethyleneimine (HEEI)					

e. E-caprolactone (wt%)

21.2

20.7

40.0

C. acid value (AV/NV)

10

15

20

25

30

35

B. viscosity (25°C Gardner)

Y<2

W-X

×

ΥX

65.1 64.8

65.2

65.0

65.l

15.0

15.1 2₁<2₂

20.2

10.0

A. non-volatile content (%)

characteristics of resin

Table 1 (continued) branched type polyester resin Synthetic Example oil-free polyester resin Composition and characteristics	olyester 6 B-©	r resin	B-8	B©	10 B-(())
oil-free polyester resin		B-(7)	в-®	в9	B-(i)
Composition and characteristics		•	,		
resin specifications					
a. number average molecular weight	1,800	1,800 2,200 3,000	3,000	1,900	2,000
b. OH value	91	64	55	93	86
c. average OH number per molecule	3.0	2.5	2.9	3.2	ა ა
d. trivalent/tetravalent alcohol (wt%) 10.4	10.4	6.2	5.2	12.5	11.7

55

Table 1 (continued)

		branched	type	polyester	resin
70					
10	Synthetic Example			11	12
	oil-free polyester res	ln		в-(1)	B-12
75	Composition and charact	eristics			
	resin raw materials				
20	1. terephthalic acid				
	2. isophthalic acid			303.9	365.6
	3. phthalic anhydride				
25	4. adipic acid			178.2	214.4
	5. azelaic acid				
30	6. sebacic acid				
	7. hexahydro phthalic	anhydride			
	8. ethyleneglycol			•	
35	9. 1,4-butanediol		•		
	10. 1,6-hexanediol			62.8	207.3
40	ll. neopentyl glycol			55.4	182.8
	12. trimethylol propand	e		399.7	30.0
	13. pentaerythritol				
45	14. Cardura E-10				
	15. &-caprolactone(n=5)			
50	16. phthalic anhydride			•	
	(for later charge)				
	17. 2-hydroxyethyl				
55	ethyleneimine (HEE	I) ·			

Table 1 (continued)

branched type polyester resin

20	Synthetic Example	11	12
	oil-free polyester resin	B-(1)	в-(12)
25	Composition and characteristics		
	resin specifications		
	a. number average molecular weight	800	4,000
30	b. OH value	315	32
	c. average OH number per molecule	4.5	2.3
35	d. trivalent/tetravalent alcohol (wt%)	39.9	3.0
	e. E-caprolactone (wt%)	0	0
	characteristics of resin		
40	A. non-volatile content (%)	65.2	65.1
	B. viscosity (25°C Gardner)	UV .	$^{\mathbf{z}}$ 1
45	C. acid value (AV/NV)	10.0	10.0

Table 1 (continued)

comparative resin

Synthetic Example	13	14	15	16	17
oil-free polyester resin	c-(13)	C-14	C-(15)	C- (6	C-(1)
Composition and characteristics					•
resin raw materials					
l. terephthalic acid	244.9		•		.
2. isophthalic acid	244.9	317.0	284.6	166.5	247.6
3. phthalic anhydride			104.9		147.2
4. adipic acid		278.9	250.4	97.6	145.2
5. azelaic acid					
6. sebacic acid	298.3				
7. hexahydro phthalic anhydride			•		-
8. ethyleneglycol	138.4	150.8	134.3		
9. 1,4-butanediol					-
10. 1,6-hexanediol					(
ll. neopentyl glycol	232.4	253.4	225.8	114.7	100.2
12. trimethylol propane		•		118.6	113.7
13. pentaerythritol	-	-			246.1
14. Cardura E-10		-	-		
15. &-caprolactone(n=5)			٠	502.7	
16. phthalic anhydride	-		-		
(for later charge)					
17. 2-hydroxyethyl					
ethyleneimine (HEEI)				•	

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Table
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cont
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continue
continued)

comparative resin

						_					_	0	"
	c.	B	Α.	lo	•	a.	Ω •	0	(o	113	Com	11.	ŝуn
·	acid va	viscosi	non-vol	haracte	E-capro	trivale	average	OH value	number	resin sp	positio	-free p	thetic
	acid value (AV/NV)	lty (25°	latile c	characteristics	E-caprolactone (wt%)	nt/tetr	OH num	Ō.	average	specifications	n∙and c	oil-free polyester	Synthetic Example
	/NV)	viscosity (25°C Gardner)	non-volatile content (%)	of resin	(wt%)	avalent a	average OH number per molecule		number average molecular weight	tions	Composition and characteristics	r resin	
			•			trivalent/tetravalent alcohol (wt%)	olecule		r weight		stics		
						œ œ							
	2.0		40.2		i	0	2:0	.6.0	20,000			C-(13)	13
			7									Ċ	
-	2.0	ΥY	70.2		Ì	0	2.0	138	800			c-(4)	14
	2.0	Z-Z	70.1		1		1.5	28	3,000			c-((5)	. 15
	ب		6		<u>5</u>	H			2,000			ç	
	10.0	×	65.1		50.3	1.9	3.5	98	000			c-(6)	16
	10.0	2 ₆ <	65.4		1	24.6	5.0	155	1,800			c-(j)	17

	_	Table 2	2		,	
Example	1	2	3	4	5	6
resin formula						
linear type polyester						
resin	A-D	A-2	A-3	A-4	A-4	A-(5)
solid	50	50	30	30	30	50
branched type polyester			-	•		l
resin	в-6	B-(7)	в-(8)	в-9	в-10	B-(8)
solid	50	50	70	70	. 70	50
OH value of blend	56	50	52	65	87	74
methylolated melamine	•					:
resin (solid)	30	30	30	30	30	30
butylated melamine (solid)						
MeMF/BuMF (solid)		•				
isocyanate (Sumidul N-75)	-					
Test results	•					
non-volatile content(%)						(
of coating composition	70.8	70.5	70.5	70.6	70.1	70.6
gloss	91.2	89.5	92.0	90.1	88.9	90.1
pencil hardness	2H	2H	2H	2Н~3Н	3H	2H
bending properties	2 T	2T~3T	3 T ·	3 T	3T	3 T
fastness to boiling water	0	0		. 0		0
chemical resistance	0	 O	0	0	0	0
marker stain resistance	\circ	0	0	0	O .	0
weather resistance	82	85	80	82	79	80

Table 2 (continued)

Example	7	8	9	10	11
resin formula					
linear type polyester					
resin	A-2	A-3	A-(1)	A-2	A-2
solid	90	80	60	40	80
branched type polyester		•			
resin	B-(1)	B-(12)	в - 6	B-(7)	B-(1)
solid	10	20	40	60	20
OH value of blend	63	42	49	52	89
methylolated melamine		·		-	Zeig (da
resin (solid)	15	35	21	15	retatives
butylated melamine (solid)			9 .	15	is
MeMF/BuMF (solid)			70/30	50/50	·
isocyanate (Sumidul N-75)					16.7
Test results					
non-volatile content(%)		. –			
of coating composition	70.2	71.2	70.1	70.2	71.3
gloss	90.2	90.5	88.6	. 88.3	90.2
pencil hardness	2Н	ЗН _	28	2H	2Н
bending properties	1T~2T	3 T	2T	3T	2 T
iastness to boiling water	0	0	0	0	0
chemical resistance	0	0	0	\circ	0
marker stain resistance	\circ	0	0	0	0
weather resistance	80	82	80	81	85
wedener reprocured	•				

5	Tabl	Table 2 (continued)		-			
3							
	Comp.Example	1	2	3	4	5	
10	resin formula						
	linear type polyester				•	•	
15	resin	c-13	C-(14)	c-(15)	A-(1)	A-4	
	solid ·	40	40	20	80	20	
	branched type polyester						(
. 20	resin	B - 9	B-8	B-8	B-6	B-10	
	solid	60	60	80	20	80	
25	OH value of blend	58	88	50	3.5	113	
	methylolated melamine	•					
	resin (solid)	30	30	30	30	30	
30	butylated melamine (solid	i)		•			
	MeMF/BuMF (solid)						
35	isocyanate (Sumidul N-75))					
	Test results			,		,	ſ
	non-volatile content(%)					•	,
40	of coating composition	55.7	70.3	70.1	70.3	70.3	
	gloss	63.2	88.4	89.2	88.3	85.3	
45	pencil hardness	. B	2H	2H	2H	2H	
	bending properties	1T	6T	3 T	2 T	6T~7T	
	fastness to boiling water	× ×	×	×	×	0	
50	chemical resistance	×	O .	X	X	\circ	
	marker stain resistance	×	\circ	×	X	0	
55	weather resistance	62	50	42	43	69	

Table 2 (continued)

	Comp. Example	6	7	8	9
10	resin formula				
	linear type polyester				
	resin	A-4	A-3		A-4
75	solid	40	100		80
	branched type polyester	•			
20	resin	c-(1)	•	в-(7)	c-(7)
	solid	60		100	20
	OH value of blend	84	45	64	81
25	methylolated melamine				
	resin (solid)	30	30	30	30
30	butylated melamine (solid)	olid)			
	MeMF/BuMF (solid)				
	isocyanate (Sumidul N-75)				
35	Test results		•		,
	non-volatile content(%)				
40	of coating composition	70.6	70.5	70.3	65.3
	gloss	87.2	80.2	88.1	*
	pencil hardness	нв	2B	3H	2H
45	bending properties	2T	lT	10T<	3 T
	fastness to boiling water		×		×
50	chemical resistance	\circ	×	0	X.
	marker stain resistance	0	×	0	×·
Δ	weather resistance	78	45	79	62
55	*loss in gloss				

Claims

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1. A coating composition for coil coating use consisting essentially of

(A) a binder component comprising a mixture of a linear polyester resin and a branched polyester resin, having a hydroxyl value of 40 to 90, said linear polyester resin being composed of polybasic acid component and polyhydric alcohol component consisting of trihydric and/or tetrahydric alcohol (0 to 10 wt% of the solid matter) and dihydric alcohol, and having 1.8 to 2.2 hydroxyl groups per molecule and a number average molecular weight of 1000 to 5000, and said branched polyester resin being composed of polybasic acid component and polyhydric alcohol component consisting of trihydric and/or tetrahydric alcohol (3 to 40 wt% of the solid matter) and dihydric alcohol, and having 2.3 to 4.5 hydroxyl groups per molecule, and

(B) a crosslinking component consisting of melamine resin, at least part of which is a methyl methylolated melamine, or isocyanate compound.

2. A composition according to claim 1 wherein the branched polyester resin is modified with a lactone of the formula:

wherein n is an integer of 4 to 10.

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EUROPEAN SEARCH REPORT

Application number

EP 86 30 6210

ategory		h indication, where appropriate, ant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.4)	
x	US-A-3 345 429 ELECTRIC CORP.) * Claims 4-6 *	(WESTINGHOUSE	1	но1 в 3/30	
A	CHEMICAL ABSTRACTION 1981, page 90, at 210367s, Columbut F.T. NOUWENS: "Findustrial coation of FATIPEC 1980, 15 II-270-II-280 * Abstract *	abstract no. 1s, Ohio, US; High solids Ings", & CONGR.	1,2		
A	CHEMICAL ABSTRAC 4, 28th July 197 abstract no. 300 Ohio, US; & JP-A (DAINIPPON INK A INC.) 09-09-1974 * Abstract *	029x, Columbus, A-74 94 723 AND CHEMICALS	1,2	TECHNICAL FIELDS SEARCHED (Int. Cl.4)	
				C 09 D C 08 L H 01 B	
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	The present search report has b	een drawn up for all claims	1		
Place of search THE HAGUE Date of completion of the search 27-02-1987		STA	Examiner STANGE		
Y: pa	CATEGORY OF CITED DOCL rticularly relevant if taken alone rticularly relevant if combined w cument of the same category chnological background	E : earlier pa after the ith another D : documer	principle undi stent documen filing date at cited in the a at cited for other	erlying the invention t, but published on, or application er reasons	